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(54) Title: FUSION PROTEINS CONTAINING N-TERMINAL FRAGMENTS OF HUMAN SERUM ALBUMIN

(57) Abstract

A fusion polypeptide comprising, as at least part of the N-terminal portion thereof, an N-terminal portion of HSA or a variant thereof and, as at least part of the C-terminal portion thereof, another polypeptide except that, when the said N-terminal portion of HSA is the 1-n portion where n is 369 to 419 or a variant thereof, then the said polypeptide is one of various specified entities, including the 585 to 1578 portion of human fibronectin or a variant thereof. The HSA-like portion may have additional N-terminal residues, such as secretion leader sequences (signal sequences). The C-terminal portion is preferably the 585-1578 portion of human plasma fibronectin. The N-terminal and C-terminal portions may be cleavable to yield the isolated C-terminal portion, with the N-terminal portion having served to facilitate secretion from the host.

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Fusion proteins containing N-terminal fragments of human serum albumin

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The present invention relates to fusion polypeptides where two individual polypeptides or parts thereof are fused to form a single amino acid chain. Such fusion may arise from the expression of a single continuous coding sequence formed by recombinant DNA techniques.

Fusion polypeptides are known, for example those where a polypeptide which is the ultimately desired product of the process is expressed with an N-terminal "leader sequence" which encourages or allows secretion of the polypeptide from the cell. An example is disclosed in EP-A-116 201 (Chiron).

Human serum albumin (HSA) is a known protein found in the blood. EP-A-147 198 (Delta Biotechnology) discloses its expression in a transformed host, in this case yeast. Our earlier application EP-A-322 094 discloses N-terminal fragments of HSA, namely those consisting of residues 1-n where n is 369 to 419, which have therapeutic utility. The application also mentions the possibility of fusing the C-terminal residue of such molecules to other, unnamed, polypeptides.

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One aspect of the present invention provides a fusion polypeptide comprising, as at least part of the N-terminal portion thereof, an N-terminal portion of HSA or a variant thereof and, as at least part of the C-terminal portion thereof, another polypeptide except that, when the said Nterminal portion of HSA is the 1-n portion where n is 369 to 419 or a variant thereof then the said polypeptide is (a) the 585 to 1578 portion of human fibronectin or a variant thereof, (b) the 1 to 368 portion of CD4 or a variant thereof, (c) platelet derived growth factor, or a variant thereof, (d) transforming growth factor, or a variant thereof, (e) the 1-261 portion of mature human plasma fibronectin or a variant thereof, (f) the 278-578 portion of mature human plasma fibronectin or a variant thereof, (g) the 1-272 portion of mature human von Willebrand's Factor or a variant thereof, or (h) alpha-1-antitrypsin or a variant thereof.

The N-terminal portion of HSA is preferably the said 1-n portion, the 1-177 portion (up to and including the cysteine), the 1-200 portion (up to but excluding the cysteine) or a portion intermediate 1-177 and 1-200.

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"human serum albumin" (HSA) is intended to include (but not necessarily to be restricted to) known or vet-to-be-discovered polymorphic forms of HSA. For example, albumin Naskapi has Lys-372 in place of Glu-372 and pro-albumin Christchurch has an altered pro-sequence. The term "variants" is intended to include (but not necessarily to be restricted to) minor artificial variations in sequence (such as molecules lacking one or a few residues, having conservative substitutions or minor insertions of residues, or having minor variations of amino acid structure). Thus polypeptides which have 80%, preferably 85%, 90%, 95% or 99%, homology with HSA are deemed to be "variants". It is also preferred for such variants to be physiologically equivalent to HSA; that is say, variants preferably share at least one to pharmacological utility with HSA. Furthermore, any putative variant which is to be used pharmacologically should be non-immunogenic in the animal (especially human) being treated.

Conservative substitutions are those where one or more amino acids are substituted for others having similar properties such that one skilled in the art of polypeptide chemistry would expect at least the secondary structure, and preferably the tertiary structure, of the polypeptide to be substantially unchanged. For example, typical such

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substitutions include asparagine for glutamine, serine for asparagine and arginine for lysine. Variants may alternatively, or as well, lack up to ten (preferably only one or two) intermediate amino acid residues (ie not at the termini of the said N-terminal portion of HSA) in comparison with the corresponding portion of natural HSA; preferably any such omissions occur in the 100 to 369 portion of the molecule (relative to mature HSA itself) (if present). Similarly, up to ten, but preferably only one or two, amino acids may be added, again in the 100 to 369 portion for preference (if present). The term "physiologically functional equivalents" also encompasses larger molecules comprising the said sequence plus a further sequence at the N-terminal (for example, pro-HSA, pre-pro-HSA and met-HSA).

Clearly, the said "another polypeptide" in the fusion compounds of the invention cannot be the remaining portion of HSA, since otherwise the whole polypeptide would be HSA, which would not then be a "fusion polypeptide".

Even when the HSA-like portion is not the said 1-n portion of HSA, it is preferred for the non-HSA portion to be one of the said (a) to (h) entities.

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The 1 to 368 portion of CD4 represents the first four disulphide-linked immunoglobulin-like domains of the human T lymphocyte CD4 protein, the gene for and amino acid sequence of which are disclosed in D. Smith et al (1987) Science 328, 1704-1707. It is used to combat HIV infections.

The sequence of human platelet-derived growth factor (PDGF) is described in Collins et al (1985) Nature 316, 748-750. Similarly, the sequence of transforming growth factors β (TGF- β) is described in Derynck et al (1985) Nature 316, 701-705. These growth factors are useful for wound-healing.

A cDNA sequence for the 1-261 portion of Fn was disclosed in EP-A-207 751 (obtained from plasmid pFH6 with endonuclease PvuII). This portion binds fibrin and can be used to direct fused compounds to blood clots.

A cDNA sequence for the 278-578 portion of Fn, which contains a collagen-binding domain, was disclosed by R.J. Owens and F.E. Baralle in 1986 E.M.B.O.J. <u>5</u>, 2825-2830. This portion will bind to platelets.

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The 1-272 portion of von Willebrand's Factor binds and stabilises factor VIII. The sequence is given in Bontham et al, Nucl. Acids Res. 14, 7125-7127.

Variants of alpha-1-antitrypsin include those disclosed by Rosenburg et al (1984) Nature 312, 77-80. In particular, the present invention includes the Pittsburgh variant (Met³⁵⁸ is mutated to Arg) and the variant where Pro³⁵⁷ and Met³⁵⁸ are mutated to alanine and arginine respectively. These compounds are useful in the treatment of septic shock and lung disorders.

Variants of the non-HSA portion of the polypeptides of the invention include variations as discussed above in relation to the HSA portion, including those with conservative amino acid substitutions, and also homologues from other species.

The fusion polypeptides of the invention may have N-terminal amino acids which extend beyond the portion corresponding to the N-terminal portion of HSA. For example, if the HSA-like portion corresponds to an N-terminal portion of mature HSA, then pre-, pro-, or pre-pro sequences may be added thereto, for example the yeast alpha-factor leader sequence. The fused leader portions of WO 90/01063 may be used. The polypeptide which is

fused to the HSA portion may be a naturally-occurring polypeptide, a fragment thereof or a novel polypeptide, including a fusion polypeptide. For example, in Example 3 below, a fragment of fibronectin is fused to the HSA portion via a 4 amino acid linker.

It has been found that the amino terminal portion of the HSA molecule is so structured as to favour particularly efficient translocation and export of the fusion compounds of the invention in eukaryotic cells.

A second aspect of the invention provides a transformed host having a nucleotide sequence so arranged as to express a fusion polypeptide as described above. By "so arranged", we mean, for example, that the nucleotide sequence is in correct reading frame with an appropriate RNA polymerase binding site and translation start sequence and is under the control of a suitable promoter. The promoter may be homologous with or heterologous to the host. Downstream (3') regulatory sequences may be included if desired, as is known. The host is preferably yeast (for example Saccharomyces spp., e.g. S. cerevisiae; Kluyveromyces spp., e.g. K. lactis; Pichia spp.; or Schizosaccharomyces spp., e.g. S. pombe) but may be any

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other suitable host such as <u>E. coli</u>, <u>B. subtilis</u>, <u>Aspergillus</u> spp., mammalian cells, plant cells or insect cells.

A third aspect of the invention provides a process for preparing a fusion polypeptide according to the first aspect of the invention by cultivation of a transformed host according to the second aspect of the invention, followed by separation of the fusion polypeptide in a useful form.

A fourth aspect of the invention provides therapeutic methods of treatment of the human or other animal body comprising administration of such a fusion polypeptide.

In the methods of the invention we are particularly concerned to improve the efficiency of secretion of useful therapeutic human proteins from yeast and have conceived the idea of fusing to amino-terminal portions of HSA those proteins which may ordinarily be only inefficiently secreted. One such protein is a potentially valuable wound-healing polypeptide representing amino acids 585 to 1578 of human fibronectin (referred to herein as Fn 585-1578). As we have described in a separate application (filed simultaneously herewith) this molecule contains cell spreading, chemotactic and chemokinetic activities

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useful in healing wounds. The fusion polypeptides of the present invention wherein the C-terminal portion is Fn 585-1578 can be used for wound healing applications as biosynthesised, especially where the hybrid human protein topically applied. However, the portion be will representing amino acids 585 to 1578 of human fibronectin can if desired be recovered from the fusion protein by preceding the first amino acid of the fibronectin portion by amino acids comprising a factor X cleavage site. After isolation of the fusion protein from culture supernatant, the desired molecule is released by factor X cleavage and purified by suitable chromatography (e.g. ion-exchange chromatography). Other sites providing for enzymatic or chemical cleavage can be provided, either by appropriate juxtaposition of the N-terminal and C-terminal portions or by the insertion therebetween of an appropriate linker.

At least some of the fusion polypeptides of the invention, especially those including the said CD4 and vWF fragments, PDGF and $\alpha_1 AT$, also have an increased half-life in the blood and therefore have advantages and therapeutic utilities themselves, namely the therapeutic utility of the non-HSA portion of the molecule. In the case of $\alpha_1 AT$ and others, the compound will normally be administered as

a one-off dose or only a few doses over a short period, rather than over a long period, and therefore the compounds are less likely to cause an immune response.

EXAMPLES : SUMMARY

Standard recombinant DNA procedures were as described by Maniatis et al (1982 and recent 2nd edition) unless otherwise stated. Construction and analysis of phage M13 recombinant clones was as described by Messing (1983) and Sanger et al (1977).

DNA sequences encoding portions of human serum albumin used in the construction of the following molecules are derived from the plasmids mHOB12 and pDBD2 (EP-A-322 094, Delta Biotechnology Ltd, relevant portions of which are reproduced below) or by synthesis of oligonucleotides equivalent to parts of this sequence. DNA sequences encoding portions of human fibronectin are derived from the plasmid pFHDEL1, or by synthesis of oligonucleotides equivalent to parts of this sequence. Plasmid pFHDEL1, which contains the complete human cDNA encoding plasma fibronectin, was obtained by ligation of DNA derived from plasmids pFH6, 16, 54, 154 and 1 (EP-A-207 751; Delta Biotechnology Ltd).

This DNA represents an mRNA variant which does not contain the 'ED' sequence and had an 89-amino acid variant of the III-CS region (R.J. Owens, A.R. Kornblihtt and F.E. Baralle (1986) Oxford Surveys on Eukaryotic Genes 3 141-160). The map of this vector is disclosed in Fig. 11 and the protein sequence of the mature polypeptide produced by expression of this cDNA is shown in Fig. 5.

Oligonucleotides were synthesised on an Applied Biosystems 380B oligonucleotide synthesiser according to the manufacturer's recommendations (Applied Biosystems, Warrington, Cheshire, UK).

An expression vector was constructed in which DNA encoding the HSA secretion signal and mature HSA up to and including the 387th amino acid, leucine, fused in frame to DNA encoding a segment of human fibronectin representing amino acids 585 to 1578 inclusive, was placed downstream hybrid promoter of EP-A-258 067 (Delta the of Biotechnology), which is a highly efficient galactoseinducible promoter functional in Saccharomyces cerevisiae. The codon for the 1578th amino acid of human fibronectin was directly followed by a stop codon (TAA) and then the S. cerevisiae phosphoglycerate kinase (PGK) gene transcription terminator. This vector was then introduced into S. cerevisiae by transformation, wherein it directed

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the expression and secretion from the cells of a hybrid molecule representing the N-terminal 387 amino acids of HSA C-terminally fused to amino acids 585 to 1578 of human fibronectin.

In a second example a similar vector is constructed so as to enable secretion by <u>S. cerevisiae</u> of a hybrid molecule representing the N-terminal 195 amino acids of HSA C-terminally fused to amino acids 585 to 1578 of human fibronectin.

Aspects of the present invention will now be described by way of example and with reference to the accompanying drawings, in which:

Figure 1 (on two sheets) depicts the amino acid sequence currently thought to be the most representative of natural HSA, with (boxed) the alternative C-termini of HSA(1-n);

Figure 2 (on two sheets) depicts the DNA sequence coding for mature HSA, wherein the sequence included in Linker 3 is underlined;

Figure 3 illustrates, diagrammatically, the construction of mHOB16;

Figure 4 illustrates, diagrammatically, the construction of pHOB31;

Figure 5 (on 6 sheets) illustrates the mature protein sequence encoded by the Fn plasmid pFHDEL1;

Figure 6 illustrates Linker 5, showing the eight constituent oligonucleotides;

Figure 7 shows schematically the construction of plasmid pDBDF2;

Figure 8 shows schematically the construction of plasmid pDBDF5;

Figure 9 shows schematically the construction of plasmid pDBDF9;

Figure 10 shows schematically the construction of plasmid DBDF12, using plasmid pFHDEL1; and

Figure 11 shows a map of plasmid pFHDEL1.

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EXAMPLE 1 : HSA 1-387 FUSED TO Fn 585-1578

The following is an account of a preparation of plasmids comprising sequences encoding a portion of HSA, as is disclosed in EP-A-322 094.

The human serum albumin coding sequence used in the construction of the following molecules is derived from the plasmid M13mp19.7 (EP-A-201 239, Delta Biotech- nology Ltd.) or by synthesis of oligonucleotides equivalent to parts of this sequence. Oligonucleotides were synthesised using phosphoramidite chemistry on an Applied Biosystems 380B oligonucleotide synthesizer according to the manufacturer's recommendations (AB Inc., Warrington, Cheshire, England).

An oligonucleotide was synthesised (Linker A) which represented a part of the known HSA coding sequence (Figure 2) from the PstI site (1235-1240, Figure 2) to the codon for valine 381 wherein that codon was changed from GTG to GTC:

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	D	P	H	E	С	Y
5′	GAT	CCT	CAT	GAA	TGC	TAT
3' ACGT	CTA	GGA	GTA	CTT	ACG	ATA
			1247			

A	K	V	F	D	E	F	K
GCC	AAA	GTG	TTC	GAT	GAA	$\mathbf{T}\mathbf{T}\mathbf{T}$	AAA
CGG	TTT	CAC	AAG	CTA	CTT	AAA	TTT
		1267					
P	L	V					

P L V
CTT GTC 3'
GGA CAG 5'

Linker 1 was ligated into the vector M13mp19 (Norrander et al, 1983) which had been digested with PstI and HincII and the ligation mixture was used to transfect E.coli strain XL1-Blue (Stratagene Cloning Systems, San Diego, CA). Recombinant clones were identified by their failure to evolve a blue colour on medium containing the chromogenic indicator X-gal (5-bromo-4-chloro-3-indolyl- β -D-galactoside) in the present of IPTG (isopropylthio- β -galactoside). DNA sequence analysis of template DNA prepared from bacteriophage particles of recombinant clones identified a molecule with the required DNA sequence, designated mHOB12 (Figure 3).

M13mp19.7 consists of the coding region of mature HSA in M13mp19 (Norrander et al, 1983) such that the codon for the first amino acid of HSA, GAT, overlaps a unique XhoI site thus:

Asp Ala

- 5' CTCGAGATGCA 3'
- 3' GAGCTCTACGT 5'

XhoI

(EP-A-210 239). M13mp19.7 was digested with XhoI and made flush-ended by S1-nuclease treatment and was then ligated with the following oligonucleotide (Linker 2):

Linker 2

5' T C T T T T A T C C A A G C T T G G A T A A A A G A 3'
3' A G A A A A T A G G T T C G A A C C T A T T T T C T 5'

<u>Hin</u>dIII

The ligation mix was then used to transfect <u>E.coli</u> XL1-Blue and template DNA was prepared from several plaques and then analysed by DNA sequencing to identify a clone, pDBD1 (Figure 4), with the correct sequence.

A 1.1 kb HindIII to PstI fragment representing the 5' end of the HSA coding region and one half of the inserted oligonucleotide linker was isolated from pDBD1 by agarose gel electrophoresis. This fragment was then ligated with double stranded mHOB12 previously digested with HindIII and PstI and the ligation mix was then used to transfect Single stranded template DNA was E.coli XL1-Blue. prepared from mature bacteriophage particles of several The DNA was made double stranded in vitro by extension from annealed sequencing primer with the Klenow fragment of DNA polymerase I in the presence deoxynucleoside triphosphates. Restriction enzyme analysis of this DNA permitted the identification of a clone with the correct configuration, mHOB15 (Figure 4).

The following oligonucleotide (Linker 3) represents from the codon for the 382nd amino acid of mature HSA (glutamate, GAA) to the codon for lysine 389 which is followed by a stop codon (TAA) and a <u>HindIII</u> site and then a <u>BamHI</u> cohesive end:

Linker 3

- E E P Q N L I K J
- 5' GAA GAG CCT CAG AAT TTA ATC AAA TAA GCTTG 3'
- 3' CTT CTC GGA GTC TTA AAT TAG TTT ATT CGAACCTAG 5'

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This was ligated into double stranded mHOB15, previously digested with <u>HincII</u> and <u>BamHI</u>. After ligation, the DNA was digested with <u>HincII</u> to destroy all non-recombinant molecules and then used to transfect <u>E.coli</u> XL1-Blue. Single stranded DNA was prepared from bacteriophage particles of a number of clones and subjected to DNA sequence analysis. One clone having the correct DNA sequence was designated mHOB16 (Figure 4).

A molecule in which the mature HSA coding region was fused to the HSA secretion signal was created by insertion of Linker 4 into <u>Bam</u>HI and <u>Xho</u>I digested Ml3mpl9.7 to form pDBD2 (Figure 4).

Linker 4

	M	K	W	V	S	F
5' GATC	C ATG	AAG	TGG	GTA	AGC	TTT
	G TAC	TTC	ACC	CAT	TCG	AAA

L I L F F L ATT TCC CTT CTT CTC **AGC** GAG TAA **AGG** GAA GAA AAA AAA TCG

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G F R V S Y S \mathbf{A} GTG GGT TTT AGG TCC GCT TAT TCG CCA CAC AAA TCC AGG CGA ATA **AGC**

R R

3' CG

GCAGCT

In this linker the codon for the fourth amino acid after the initial methionine, ACC for threonine in the HSA prepro leader sequence (Lawn et al, 1981), has been changed to AGC for serine to create a HindIII site.

A sequence of synthetic DNA representing a part of the known HSA coding sequence (Lawn et al., 1981) (amino acids 387, Fig. 2), fused to part of the known 382 fibronectin coding sequence (Kornblihtt et al., 1985) (amino acids 585 to 640, Fig. 2), was prepared by synthesising six oligonucleotides (Linker 5, Fig. 6). The oligonucleotides 2, 3, 4, 6, 7 and 8 were phosphorylated the then polynucleotide kinase and T4using oligonucleotides were annealed under standard conditions in pairs, i.e. 1+8, 2+7, 3+6 and 4+5. The annealed oligonucleotides were then mixed together and ligated with mHOB12 which had previously been digested with the The ligation restriction enzymes <u>HincII</u> and <u>EcoRI</u>.

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mixture was then used to transfect <u>E.coli</u> XL1-Blue (Stratagene Cloning Systems, San Diego, CA). Single stranded template DNA was then prepared from mature bacteriophage particles derived from several independent plaques and then was analysed by DNA sequencing. A clone in which a linker of the expected sequence had been correctly inserted into the vector was designated pDBDF1 (Fig. 7). This plasmid was then digested with <u>PstI</u> and <u>EcoRI</u> and the approx. 0.24kb fragment was purified and then ligated with the 1.29kb <u>BamHI-PstI</u> fragment of pDBD2 (Fig. 7) and <u>BamHI + EcoRI</u> digested pUC19 (Yanisch-Perron, et al., 1985) to form pDBDF2 (Fig. 7).

A plasmid containing a DNA sequence encoding full length human fibronectin, pFHDEL1, was digested with EcoRI and a 0.77kb EcoRI-XhoI fragment (Fig. 8) was isolated and then ligated with EcoRI and SalI digested M13 mp18 (Norrander et al., 1983) to form pDBDF3 (Fig. 8).

The following oligonucleotide linker (Linker 6) was synthesised, representing from the PstI site at 4784-4791 of the fibronectin sequence of EP-A-207 751 to the codon for tyrosine 1578 (Fig. 5) which is followed by a stop codon (TAA), a HindIII site and then a BamHI cohesive end:

Linker 6

G P D Q T E M T I E G L

GGT CCA GAT CAA ACA GAA ATG ACT ATT GAA GGC TTG

A CGT CCA GGT CTA GTT TGT CTT TAC TGA TAA CTT CCG AAC

Q P T V E Y Stop

CAG CCC ACA GTG GAG TAT TAA GCTTG

GTC GGG TGT CAC CTC ATA ATT CGAACCTAG

This linker was then ligated with PstI and HindIII digested pDBDF3 to form pDBDF4 (Fig. 8). The following DNA fragments were then ligated together with BglII digested pKV50 (EP-A-258 067) as shown in Fig. 8: 0.68kb ECORI-BamHI fragment of pDBDF4, 1.5kb BamHI-StuI fragment of pDBDF2 and the 2.2kb StuI-EcoRI fragment of pFHDEL1. The resultant plasmid pDBDF5 (Fig. 8) includes promoter of EP-A-258 067 to direct the expression of the HSA secretion signal fused to DNA encoding amino acids 1-387 of mature HSA, in turn fused directly and in frame human acids 585-1578 of with DNA encoding amino fibronectin, after which translation would terminate at This is then followed by the the stop codon TAA. S.cerevisiae PGK gene transcription terminator. The

plasmid also contains sequences which permit selection and maintenance in <u>Escherichia coli</u> and <u>S.cerevisiae</u> (EP-A-258 067).

This plasmid was introduced into <u>S.cerevisiae</u> S150-2B (<u>leu2-3 leu2-112 ura3-52 trp1-289 his3- 1</u>) by standard procedures (Beggs, 1978). Transformants were subsequently analysed and found to produce the HSA-fibronectin fusion protein.

EXAMPLE 2: HSA 1-195 FUSED TO Fn 585-1578

In this second example the first domain of human serum albumin (amino acids 1-195) is fused to amino acids 585-1578 of human fibronectin.

The plasmid pDBD2 was digested with <u>BamHI</u> and <u>BglII</u> and the 0.79kb fragment was purified and then ligated with <u>BamHI</u>-digested M13mp19 to form pDBDF6 (Fig. 6). The following oligonucleotide:

5'-C C A A A G C T C G A G G A A C T T C G-3'

was used as a mutagenic primer to create a XhoI site in pDBDF6 by in vitro mutagenesis using a kit supplied by Amersham International PLC. This site was created by

changing base number 696 of HSA from a T to a G (Fig. 2). The plasmid thus formed was designated pDBDF7 (Fig. 9). The following linker was then synthesised to represent from this newly created XhoI site to the codon for lysine 195 of HSA (AAA) and then from the codon for isoleucine 585 of fibronectin to the ends of oligonucleotides 1 and 8 shown in Fig. 6.

Linker 7

D E L R D E G K A S. S A K

TC GAT GAA CTT CGG GAT GAA GGG AAG GCT TCG TCT GCC AAA

A CTT GAA GCC CTA CTT CCC TTC CGA AGC AGA CGG TTT

I T E T P S Q P N S H

ATC ACT GAG ACT CCG AGT CAG C

TAG TGA CTC TGA GGC TCA GTC GGG TTG AGG GTG G

This linker was ligated with the annealed oligonucleotides shown in Fig. 3, i.e. 2+7, 3+6 and 4+5 together with XhoI and EcoRI digested pDBDF7 to form pDBDF8 (Fig. 9). Note that in order to recreate the original HSA DNA sequence, and hence amino acid sequence, insertion of linker 7 and the other oligonucleotides into pDBDF7 does not recreate the XhoI site.

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The 0.83kb <u>BamHI-StuI</u> fragment of pDBDF8 was purified and then was ligated with the 0.68kb <u>EcoRI-BamHI</u> fragment of pDBDF2 and the 2.22kb <u>StuI-EcoRI</u> fragment of pFHDEL1 into <u>BglII-digested pKV50</u> to form pDBDF9 (Fig. 9). This plasmid is similar to pDBDF5 except that it specifies only residues 1-195 of HSA rather than 1-387 as in pDBDF5.

When introduced into <u>S.cerevisiae</u> S150-2B as above, the plasmid directed the expression and secretion of a hybrid molecule composed of residues 1-195 of HSA fused to residues 585-1578 of fibronectin.

EXAMPLE 3: HSA 1-387 FUSED TO Fn 585-1578, AS CLEAVABLE MOLECULE

In order to facilitate production of large amounts of residues 585-1578 of fibronectin, a construct was made in which DNA encoding residues 1-387 of HSA was separated from DNA encoding residues 585-1578 of fibronectin by the sequence

I E G R
ATT GAA GGT AGA
TAA CTT CCA TCT

which specifies the cleavage recognition site for the blood clotting Factor X. Consequently the purified secreted product can be treated with Factor X and then the fibronectin part of the molecule can be separated from the HSA part.

To do this two oligonucleotides were synthesised and then annealed to form Linker 8.

Linker 8

G ' r I E Q N E E GGT TTA ATT GAA CAG TAA GAA GAG CCT AAT CTT CCA TTATAA GTC CTT CTC GGA

S P Q E I R CCG AGT CAG ACT GAG ATC ACT AGA TCA GTC GGG GGC CTC TGA TCT TAG TGA

N S H

TTG AGG GTG G

This linker was then ligated with the annealed oligonucleotides shown in Fig. 6, i.e. 2+7, 3+6 and 4+5 into <u>Hincil.nic.l.</u>

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(Fig. 7). The plasmid was then digested with PstI and EcoRI and the roughly 0.24kb fragment was purified and then ligated with the 1.29kb BamHI-PstI fragment of pDBD2 and BamHI and EcoRI digested pUC19 to form pDBDF11 (Fig. 10).

The 1.5kb BamHI-StuI fragment of pDBDF11 was then ligated with the 0.68kb EcoRI-BamH1 fragment of pDBDF4 and the 2.22kb StuI-EcoRI fragment of pFHDEL1 into BglII-digested pKV50 to form pDBDF12 (Fig. 10). This plasmid was then introduced into S.cerevisiae S150-2B. The purified secreted fusion protein was treated with Factor X to liberate the fibronectin fragment representing residues 585-1578 of the native molecule.

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CLAIMS

A fusion polypeptide comprising, as at least part of the N-terminal portion thereof, an N-terminal portion of HSA or a variant thereof and, as at least part of the C-terminal portion thereof, another polypeptide except that, when the said N-terminal portion of HSA is the 1-n portion where n is 369 to 419 or a variant thereof then the said polypeptide is (a) the 585 to 1578 portion of human fibronectin or a variant thereof, (b) the 1 to 368 portion of CD4 or a variant thereof, (c) platelet derived growth factor or a variant thereof, (d) transforming growth factor β or a variant thereof, (e) the 1-261 portion of mature human plasma fibronectin or a variant thereof, (f) 278-578 portion of mature human plasma the fibronectin or a variant thereof, (g) the 1-272portion of mature human von Willebrand's Factor or a variant thereof, or (h) alpha-1-antitrypsin or a variant thereof.

- 2. A fusion polypeptide according to Claim 1 additionally comprising at least one N-terminal amino acid extending beyond the portion corresponding to the N-terminal portion of HSA.
- 3. A fusion polypeptide according to Claim 1 or 2 wherein there is a cleavable region at the junction of the said N-terminal or C-terminal portions.
- 4. A fusion polypeptide according to any one of the preceding claims wherein the said C-terminal portion is the 585 to 1578 portion of human plasma fibronectin or a variant thereof.
- 5. A transformed or transfected host having a nucleotide sequence so arranged as to express a fusion polypeptide according to any one of the preceding claims.
- 6. A process for preparing a fusion polypeptide by cultivation of a host according to Claim 5, followed by separation of the fusion polypeptide in a useful form.
- 7. A fusion polypeptide according to any one of Claims 1 to 4 for use in therapy.

FIGURE 1

дsр	Ala	His	Lys	Ser	Glu	Val	Ala	His	10 Arg		Lys	ysb	Leu	Gly	Glu	: Glu	: Ast	: Phe	20 Lys
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Tyr	Lys	Ala	Ala	Phe	mh	Glu	Cys	Cys	170 Gln	Ala	Ala	yeb	Lys	Ala	Ala	Cys	Leu	Leu	180 Pro
Lys	Leu	Asp	Glu	Leu	Arg	Asp	Glu	Gly	190 Lys	Ala	Ser	Ser	Ala	Lys	Gln	Arg	Leu	Lys	200 Cys
Ala	Ser	Leu	Gln	Lys	Phe	Gly	Glu	Arg.	210 Ala	Phe	Lys	Ala	فتت	Ala	Val	Ala	AIG	Leu	220 Ser
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FIGURE 2 DNA sequence coding for mature HSA

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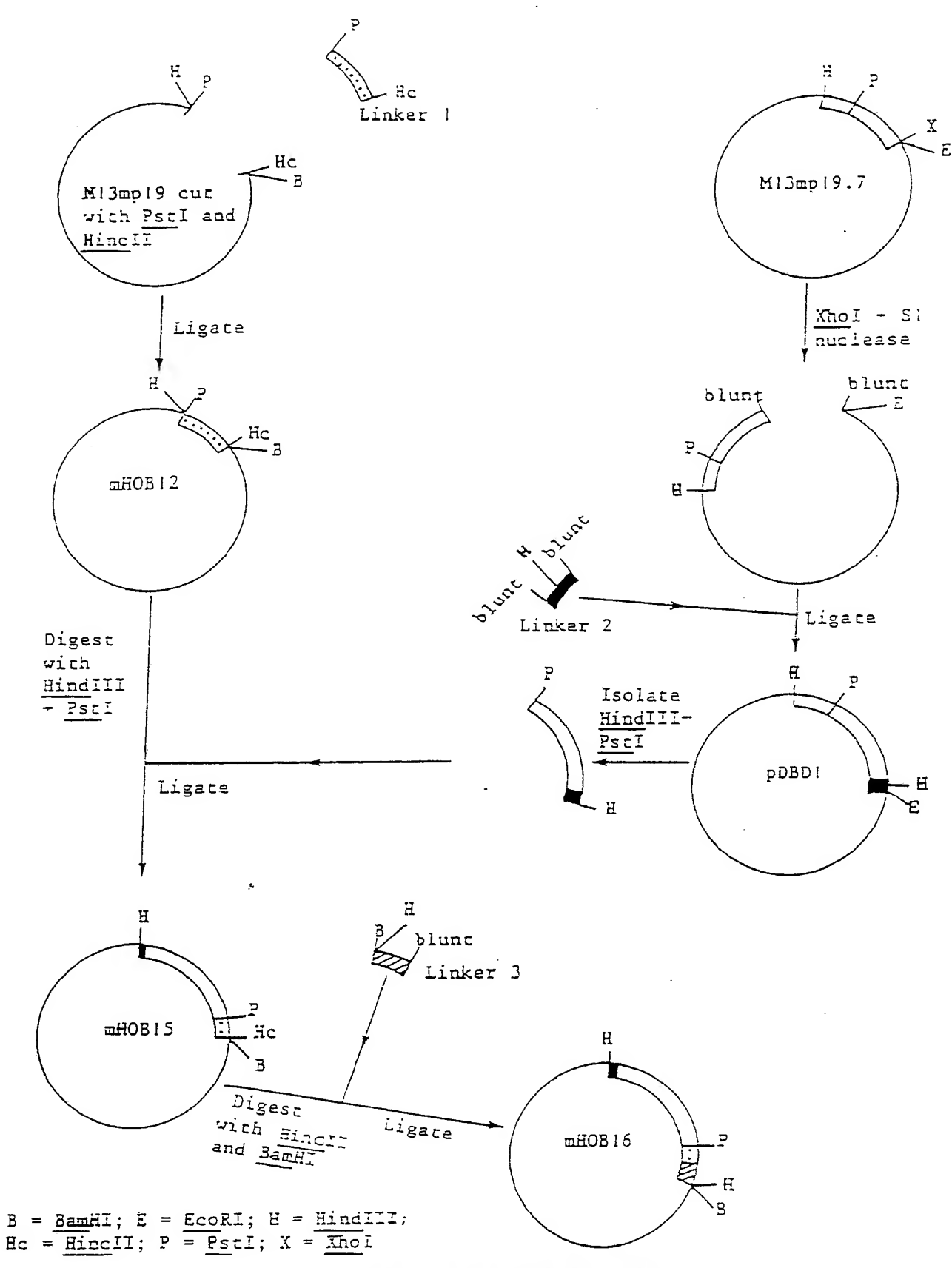
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FIGURE 2 Cont. 1050 1060 1070 1080 1090 1100 1110 1120 GAGACTTGCCAAGACATATGAAACCACTCTAGAGAAGTGCTGTGCCGCTGCAGATCCTCATGAATGCTATGCCAAAAGTGT RLAKTYETTLEKCCAAADPHECYAKV 1130 1140 1150 1160 1170 1180 1190 1200 F D E F K P L V E E P Q N L I K Q N C E L F E Ç L G E 1210 1220 1230 1240 1250 1260 1270 1280 TACAAATTCCAGAATGCGCTATTAGTTCGTTACACCAAGAAAGTACCCCAAGTGTCAACTCCAACTCTTGTAGAGGTCTC Y K F Q N A L L V R Y T K K V P Q V S T P T L V E V S 1290 1300 1310 1320 1330 1340 1350 1360 RNLGKVGSKCCKHPEAKRMPCAEDYL 1370 1380 1390 1400 1410 1420 1430 CCGTGGTCCTGAACCAGTTATGTGTGTTGCATGAGAAAACGCCAGTAAGTGACAGAGTCACAAAATGCTGCACAGAGTCC 5 V V L N Q L C V L H E K T P V S D R V T K C C T E S 1450 1460 1470 1480 1490 1500 1510 1520 TTGGTGAACAGGCGACCATGCTTTCAGCTCTGGAAGTCGATGAAACATACGTTCCCAAAGAGTTTAATGCTGAAACATT LVNRRPCFSALEVDETYVPKEFNAETF 1530 1540 1550 1560 1570 1580 1590 1600 TFHADICTLSEKERQIKKQTALVELV 1630 1640 1650 1660 1670 1680 1610 1620 AACACAAGCCCAAGGCAACAAAAGAGCAACTGAAAGCTGTTATGGATGATTTCGCAGCTTTTGTAGAGAAGTGCTGCAAG K H K P K A T K E Q L K A V M D D F A A F V E K C C K 1710 1720 1730 1740 1750 1760 1690 1700 GCTGACGATAAGGAGACCTGCTTTGCCGAGGAGGGTAAAAAACTTGTTGCTGCAAGTCAAGCTGCCTTAGGCTTATAACA

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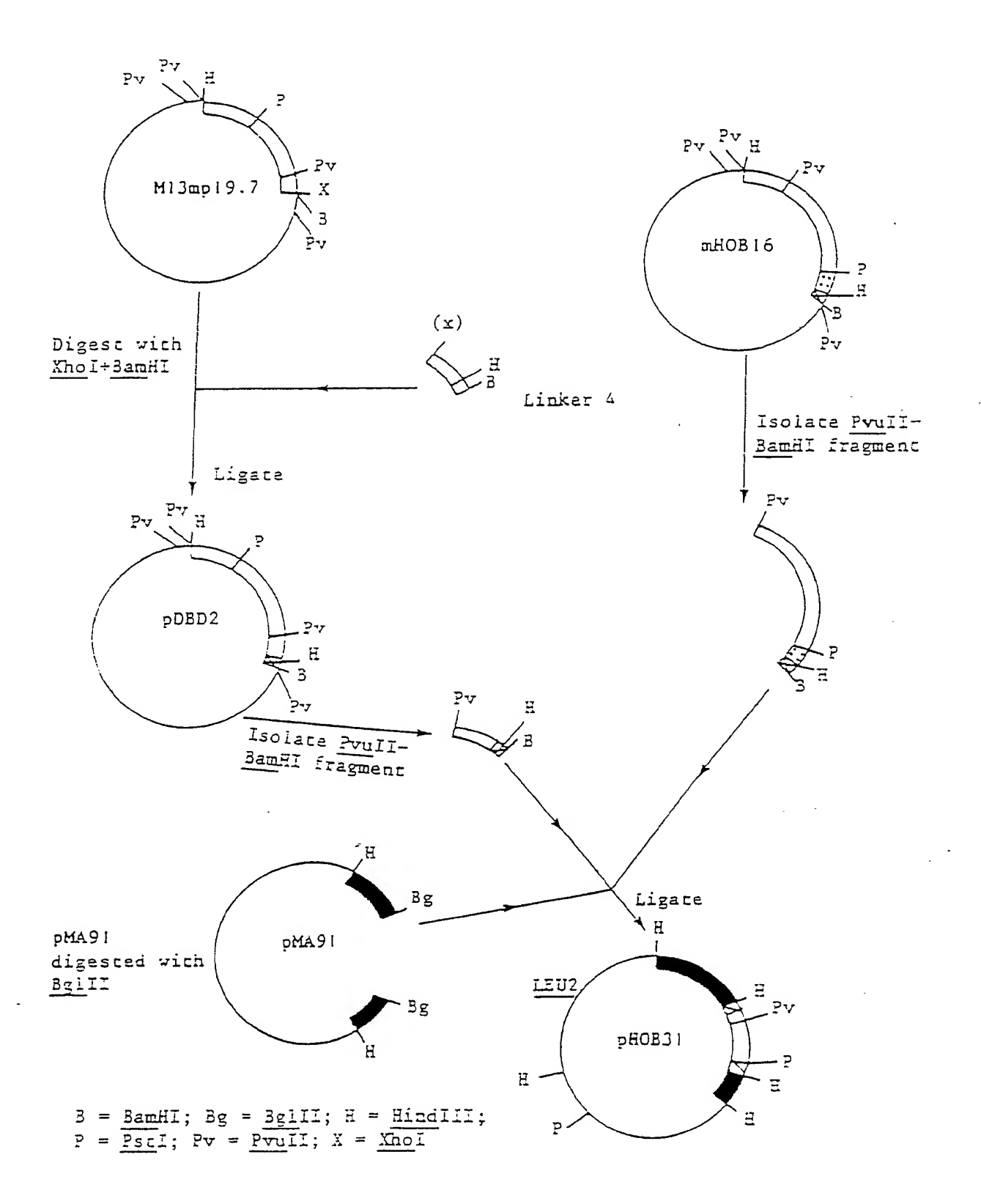
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TCTACATTTAAAAGCATCTCAG

FIGURE 3 Construction of mHOB16



SUBSTITUTE SHEET

FIGURE 4 Construction of pHOB31



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Fig. 5A

280 Asp 300 Met 360 Asn 340 Phe 200 Cys 220 Asn 0.00 0.10 0.10 260 Ala 320 Tyr 280 Val Asp Leu Met Cys CID CIL Asn Ala 中于 Thr Gln Met Lys Lys Asn Lys Lys Cys Lys Gin Arg His Gly GIn His Val Tyr Ser Gly Val Arp d p Asp Arg Asp Lys Cys Asn Asn <u>の</u>に Tyr Thr Ala 上 Ser Val Gly Val Kal Leu Arg Asn Cys Phe Gla HIS Gly Asn GIn Arg Cys Ser TH GIY Gly Ser Ser $\frac{\Omega}{\sqrt{\lambda}}$ Ser Val Gin Asn alu Gin Glu Cys Tyr 110 Cys P P Ala TY. Gin Gly Cys 174 Gly Tyr Val Ser Leu Phe 4TP Leu Lys Arg Thr Cys Pro 水 Cys Thr Tyr Thr 十九十 보 GIY Val Phe <u>S</u> pro Glu Leu Asn Asp Pro Lys Phe His Lys Thr Gly Met Thr Gly Gln AIO Gly Gly Leu Pro Asn Asp Cys <u>S</u> Gly GIn T to T Gin Tyr Ala Lys Pro Pro 116 \ \ \ \ \ \ \ Gly His Ser Phe Asn Pro Pro Trp Cys HIS Leu Arg Gly <u>aly</u> Gly りに Gly Ser 350 Asp 6 6 √ 5 √ 230 TH 250 Ser 210 Arg 290 Gin 870 61n 350 567 Lys **Met** Cys Pro Phe Lys Ser Cys חוֹם GIn GIY Thr 7 Ser Cys GIN Val Asp Arg Met Asn Arg J G Ser S Q Ŋ Gly Asn Gly Ö Ser Gln Pro 卜 7 $\overline{\sigma}$ グ $\overline{\mathfrak{Q}}$ ゴ q <u>n</u> Pro Ala Phe Ser His Thr Thr 水 <u>G</u> Cys Cys Ala Gly Val Bla His 十十十 Glu Gly GIY Trp הים Pro Arg Ser 同义 GIn Asp Gin Lys Cys Lys. Asp Asn Leu Leu Gln <u>n</u>n Lys II O <u>Val</u> Gly G S Sin Th Leu Gly Met Val Leu Pro Asn <u>Glu</u> Asp GIn Asp Thr ヤヤ Asn 보 Cys Arg Thr Thr Val Ser Val Ser 뉴 GIY ה Pro Tyr Ser ナガ Asn <u>ה</u> Gin Ala Gin Gin Cys Thr Cys Val Val Cys Asn Ser Arg Gly Lys Gly **√a**l Asp Arg Cys d d 上上 Asn ren Ser Tyr Ala Glu 中 Cys Ser Gly <u>ה</u> Thr GIY Gly Val Ely Thr cys Cys Ser Asp Tyr His Ile <u>8</u> Asn Asn Arg Gly Asp Ser Gly Thr Tyr Gly Arg

Fig. 5E

700 11e 640 Leu 666 G-12 680 Val 720 Arg 740 Thr 820 GIY **620** Val Leu Glu Phe Gly <u>G</u> Ser Tyr Thr Gly TY 늄 Sal Asp Asn Asn Gly Gin Asn Lys His 中上 Asp Asp Phe スト Tr Cys Val Gly Se Leu Ser Thr Se שׁב His Cys Asn Ile Lys Ile GIY Val Val Ser Gin Thr Ser Val 7 Asp Leu <u>G</u> Pro Pro Thr Glu Gln Thr Tyr Val Cys Ser Gly Ser 九十 Gly <u>k</u> Leu Thr Arg Gin <u>ه</u> Thr Phe Ser Asn Fro Ser Gin His Ser Lys 上げ Tyr HIS Val Asp Tyr Lys pro Thr Asp Cys $\frac{\omega}{n}$ Thr Tyr Val Ala Ser Ser מוש Ser <u>G</u>Lu Tyr Tyr Tyr Asp Ser Arg Asn Ser Gin Arg Thr Asn Ile Ala Arg Thr Thr 九 Thr Ser Gly Asp Asn Leu Leu Ala Cys Asp Ser GIn Ser S n Val Gln Ser <u>8</u> Ash Thr Asp Gin His Asp Gly Leu Val Gly 730 Asp 690 Leu 650 Leu 670 Ser 730 Val 137 Asp Glu Trp Led Gly Gin Lys 古ず Pro G G Gin <u>n</u> ∑a Va Pro Asp 古 Asp Ile Thr Gl N N aly Ser ā a p <u>></u> ġ <u>8</u> A Ś Ŋ 工 Q Phe Pro Leu Asn Leu <u>ה</u> Thr Glu Thr Ile מפ 中 Phe <u>alu</u> Val Lys His Cys Cys 水 Gly Val Pro Pro Phe Pro Trp Lys Gln שמ Va. Ser Gly 古古 His Ale Pro 첫 Met 규 <u>ว</u> Asp Asn Phe Asp Ala Arg Phe Asp Trp Glu Leu Asp <u>0</u> Gln Asp Ser Thr Val Ile Glu <u>ာ</u>ဗ Cys Val פות <u>ว</u> Pro Pro $\frac{\Omega}{r}$ Phe Va! Gln Pro Lys 十十十 Ser <u>اھ</u> Arg Ala Asp Cys Ser Arg HIS <u>aly</u> Gly Pro Trp Ser Gly שמ Thr Leu Ser Asp Glu Tyr Ser Met Lys Arg Val Gla Ala Arg Pro Asp Pro 开 Thr 10 Ser Gln GIY Met Trp Giy Ala GIY Lys Val 却 Val Lys Gly Κα

Fig. 50

200 Ser 220 Lys 180 Thr 040 Gly 1020 Tyr 060 11e 000 V&1 980 Ser 989 Pro 900 Ala Pro Glu Lys Ser ছ Asp 부 Lys Thr Ser Ala <u>S</u> HIS エド ছ Pro <u>an</u> Arg Lys Pro Lec Ile Thr 7 Gly Thr Ser Pro Leu Ser Glu Asp Pro Ala Asn ħ Pro <u>alc</u> Thr Ala S S 보 GIS <u>₩</u> Glu Gly 뉴 Ser Asp Trp Ala Val Arg Val Asn Ala Val Pro Val Leu Ser \<u>a</u> 7 Lys Pro 8 上れ His Ser Glu Pro Leu Gly <u>an</u> Thr Tyr Asn Pro Leu Val GIY 나다 Ser al S Pro Asn Val Pro Pro Leu Val Phe Val Gly Ser Gly Ile Th Asp Asn GIY 보 Arg Asn Gly Val Ala Ala 두 Gh Val Val Val Arg Ser Val Va Va Thr Met S C C Tyr Asp Gin Leu コラ 보 Asn 九十 Gh S S S ट् Sa Sa Va Va ā Glu Phe Asp Ser Arg Gly Thr Asn Ala Tyr Ţ 1230 Asp 210 Leu 190 Leu 170 Pro 090 Pro 990 Arg 110 Ser 150 His 010 010 030 Glu 050 Val 070 970 Thr Thr Ser 890 Val 910 Val Pro Ser GIY Ser Arg Gin Pro Pro Asp S S S Val Ser Asp 上上 Val Asn 0 Gly O Ash Ala Val Pro Pro Sal Asn Ala G S Th 十十十 Va Va Ala 九 q Ala Thr Pro Ser GIN Val Leu Arg Asp Thr 보 Ser 10 Asp Ser Gly Ser Ile Thr Asn Gly Gin Gly Gln Gly Pro Phe Arg Thr Glu Val Val Asn nan Val Pro Leu Ser Trp Glu Arg Leu Trp Gin Pro Lys GIC Lys Val r Olo Gly Asn Ser Tyr Asn Ser 九十 Pro Arg Arg Phe Lys Leu Asn Thr Lys <u>G</u> Val Ile GIN Tyr Gly Asp Phe <u>G</u> Ser Arg Ser Phe Asn Thr Leu Val Phe gin Pro Thr Phe Lys Leu Arg Leu Asp Gly Tyr Leu Thr Arg <u>ว</u>เย. الح <u>U</u>U Thr ren ₹ Ş Ile Val Thr Asp Val Ile Ile Ser Val ren Leu Gln Pro 环 GIN Ala Arg Val Asp Thr Arg Gly Asn

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Gln Thr

Asp

660 GIV 560 Ser 1580 Ser 1600 177 1640 Lys 620 GIn ABA <u>ದ</u> ಶ 九十 Val ヹ Seg Leu **Lys** Ser Val Va | Val Lec Thr Ala Val Y Z Asp Thr <u>つ</u> Thr Ser Val GIn Lys Sal Ser <u>8</u> Leu Pro Arg Th Val Ser Leu 古ず Ala Val Ser Pro Gln Arg Val ٦ً Ø <u>G</u> Leu Pro Gin T Š .590 Ser 670 Ser Glu leu <u>い</u> Val Gly Lle S Ası GIL GIL Ž Pro Leu Gin 10 Thr Ser tyr Lys Glu Pro Thr Asp Glu Met Pro Asn Val Lys Asn エド Pro Gln Met

Ala

Val

Met

FNDEL

Pro

aly

Pro

Th

Ala

Pro

Ala

Tyr

Val

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Arg

Pro

Pro

Ser

<u>Ray</u>

Asn

上

Val Thr

Val

<u>G</u>

2000 Thr 2060 Tyr 2020 Leu 2040 Asn 900 Pro 1980 Ser 1840 Thr 880 Pro 1920 GIY 1940 171 960 Ala 1860 Lys 800 Val 820 G/y Ash Ala Gin Leu Leu HIS Thr Lys Gin Ser Ser Ę Pro ユー Ser Sal 9 Asp Asn Asn Lys Arg <u>つ</u> Arg Cys Thr つ で Leu Thr 九十 Ser Ser Ala Lys Ile ゴド Τζ Asn <u>ต</u> Arg Pro H.s Asn Gly 上 ASP GIY Va Sa Glu Val Gin Asp Ala Val Tyr Ser Pro Phe Pro Cys Leu Lys Leu Thr Lys Pro Gly <u>Glu</u> <u>8</u> Ile Asn Thr U U Ile Tyr 110 Trp Len Lys Leu GI Thr Ile Val Va Va Tyr 五元 Pro Pro Gly Gly Lys 文 Ser Val Thr Arg Asn Phe Arg Pro r E Pro His Pro Thr Arg Thr Ser Thr Ala Pro Ile Yal 中 Val Phe Asp Aso Pro Leu <u>S</u> Ile Pro Arg 水 Ser Val Gly Gin Ile Gly Gin Ile Th GIN Phe Pro S S <u>Ka</u> Leu Gly Pro GIT 上げ Ser S S S Val Ser Ser Ala Ser S S S Tyr Val Leu cys Asp Asn Phe い に い に い Phe Asp Aso <u>つ</u> Asn Ala Arg Arg Tall Jall Gly Ser Tyr Gly 2050 四口 1890 Leu 1930 I le 990 Pro 2030 2010 1910 GIZ 1950 His 970 Pro 850 11e 750 Thr 1970 Thr 710 7hr Arg 790 Phe 1830 Pro Ala 180 171 Asp Arg Arg Met Met Thr Arg Arg Arg Lys 5 Ala Ala Arg lle 五十 Lau Asp GIn GIn מות Asp Val Phe Pro Thr Arg J. D Asn Arg Lys Glu Ile Thr Va J Thr 7 규 Asn Asp Leu Lys His Asp Ala Ser Glu Pro 水 Gla Arg Pro 十七 Ala Val Pro Val Gly Thr HIS Pro <u>n</u> Val Gly Trp Gly Asn <u>He</u> Asp Ser Arg Gly פות Thr Thr Ser GIY GIY Val <u>GI</u> Th GIn Arg Pro Thr $\frac{\partial}{\partial z}$ ナド Ser His Leu 11e Pro Pro Gly Ala Arg Tan alu Gly Val Asn Leu His Ser Asp Leu Pro 九十 Pro 규 Ala GIS Ala Arg Glu Thr <u>Val</u> Leu Pro Thr Gin Asp Pro Thr Asn GIn Asp שנת Asp UU Pro Phe Phe Thr Gly 1×r Ala ren Pro Asp <u>G</u>1 His Ala שוב Val Pro Ser Gly Pro ren Trp Pro Val Ser Pro Ile Gly Phe Leu Cys ren Ser Lys Ala His Ser Ser Gly Val

2180 Arg 2200 Gin 2160 Ala Gin Met Met Arg Pro. Ser Leu **Z**ut Cys Ale 卜 Asn Phe Tyr <u>n</u> Asp Asn Glu His ۵ľy Cys Asn Pro Cys Lys <u>D</u> 文 Asp <u>a</u> <u>a</u> Arg Ser Ile Trp Cys Trp <u>n</u> Gly Pro GIn Lys GIn ςλs Gly G_y Arg GIU Asn 2130 Glu Gly Gly Gly Gly 2210 Asn Val Gly Ser Val Asp 十十 Lys His <u>alu</u> Gly I Asn Tyr Lys' Ile Gly Glu Ls Thr Cys Leu Gly Asn Gly L r Asp Asp Gly Lys Thr Tyr F cys Ser Cys Thr Cys Phe G Gly Gly Glu Pro Ser Pro G Tyr His Gln Arg Thr Asn T r Cys Leu Gly Asn Gly L p Asp Gly Lys Thr Tyr F s Ser Cys Thr Cys Phe G f Gly Glu Pro Ser Pro G r His Gln Arg Thr Asn 7 I Gln Ala Asp Arg Glu F Val Pro Arg Asp

FIG. SF

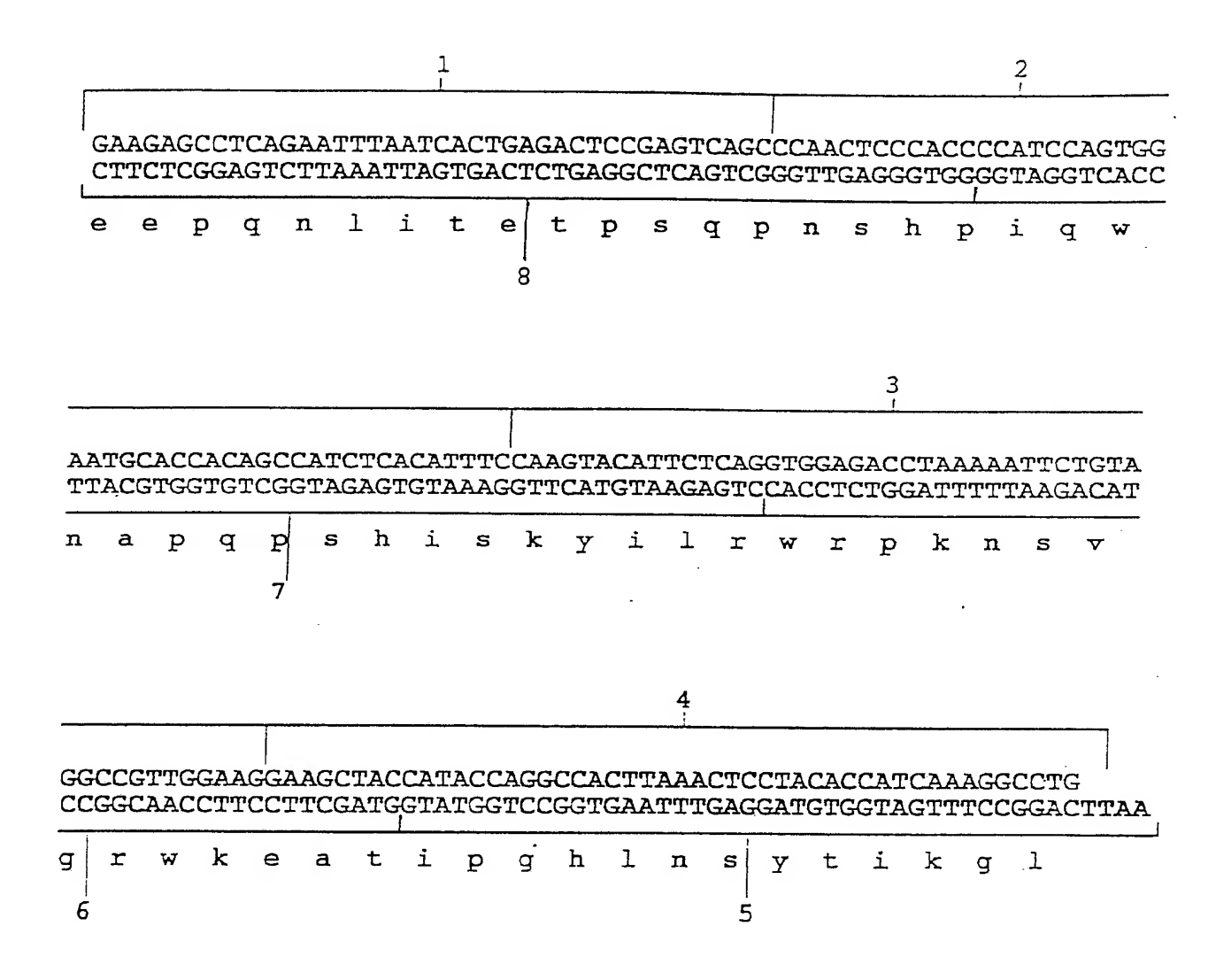


Figure 6 Linker 5 showing the eight constituent oligonucleotides

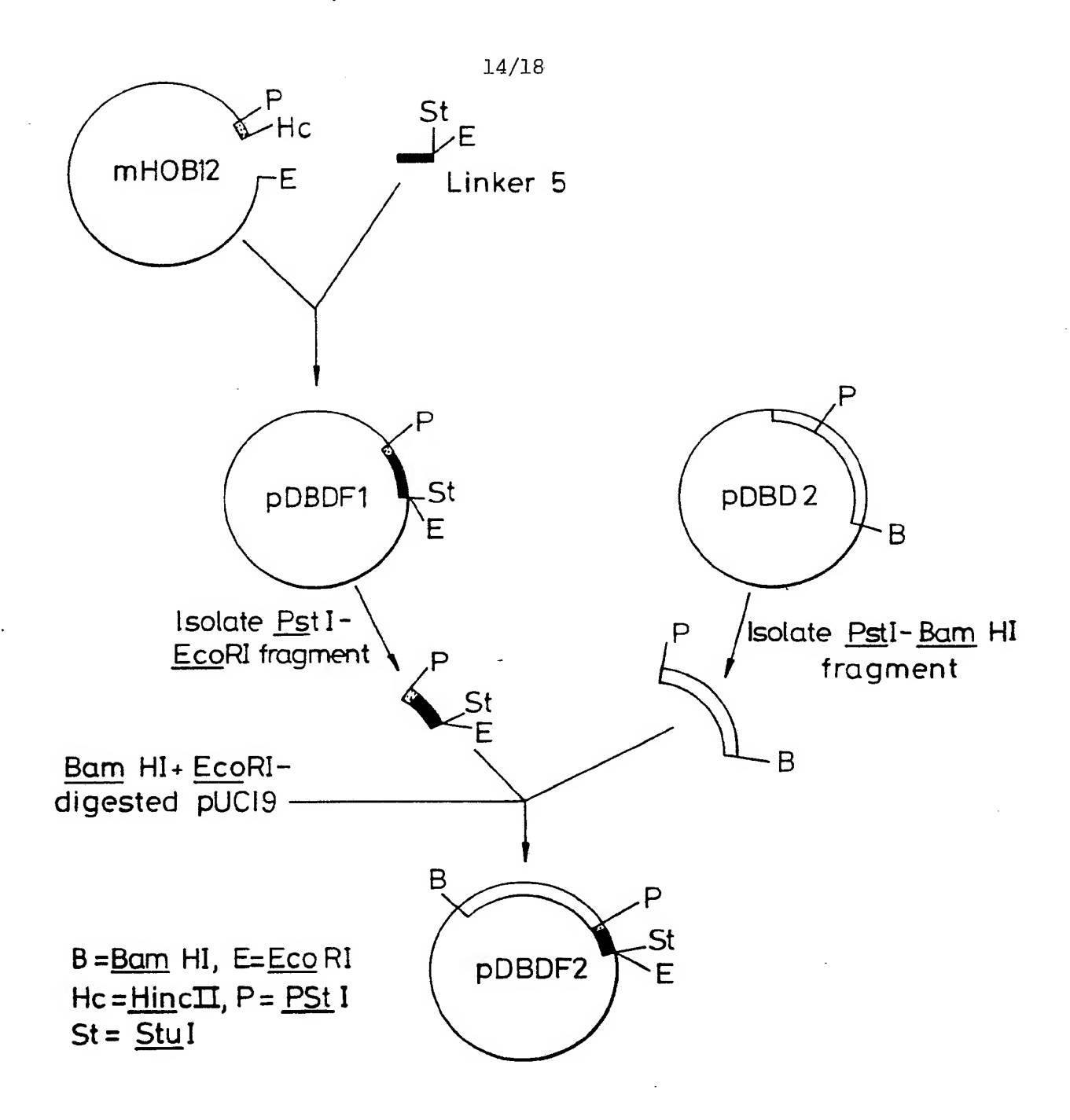


Fig. 7 Construction of pDBDF2

PCT/GB90/00650

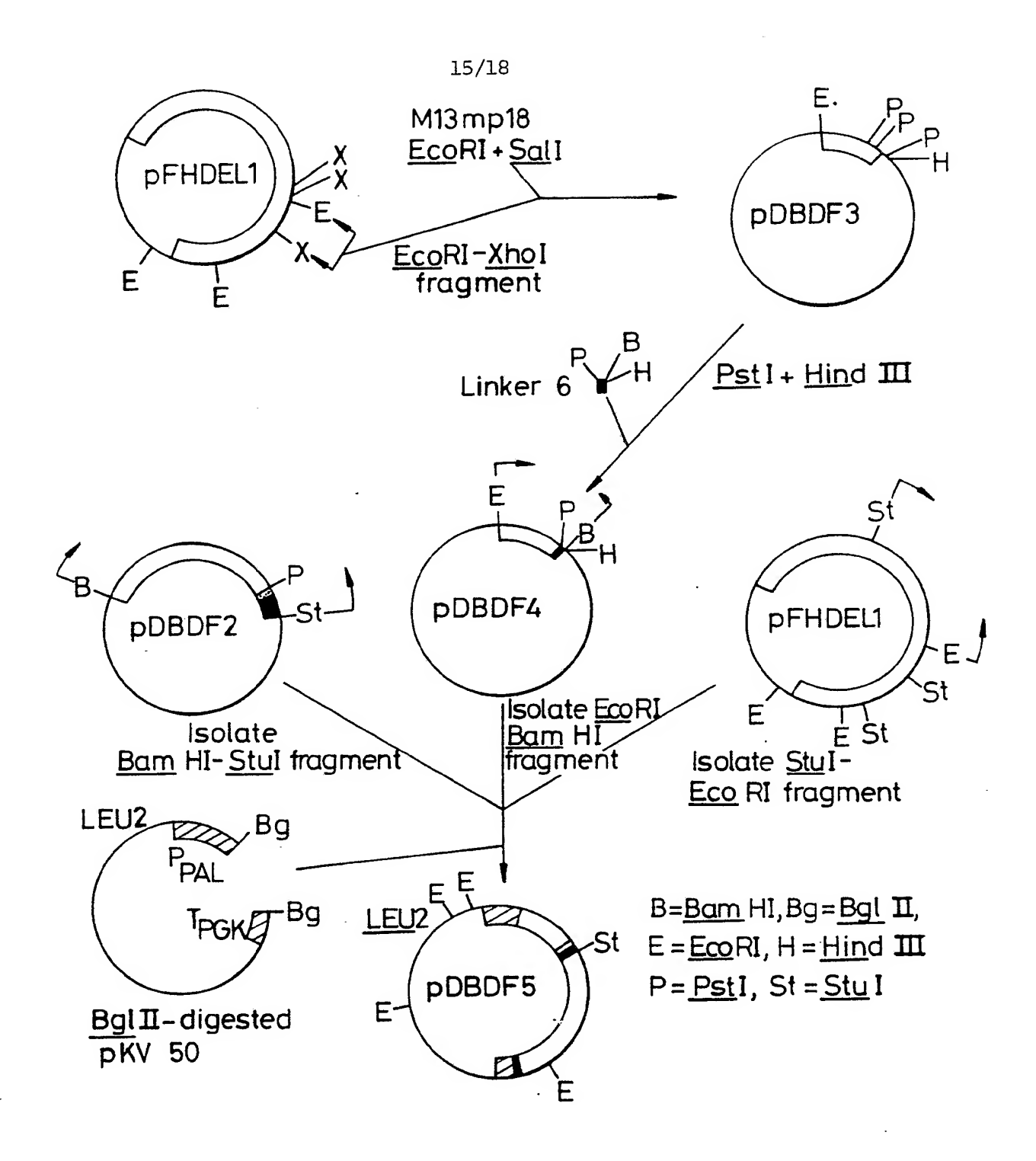


Fig. 8 Construction of pDBDF5

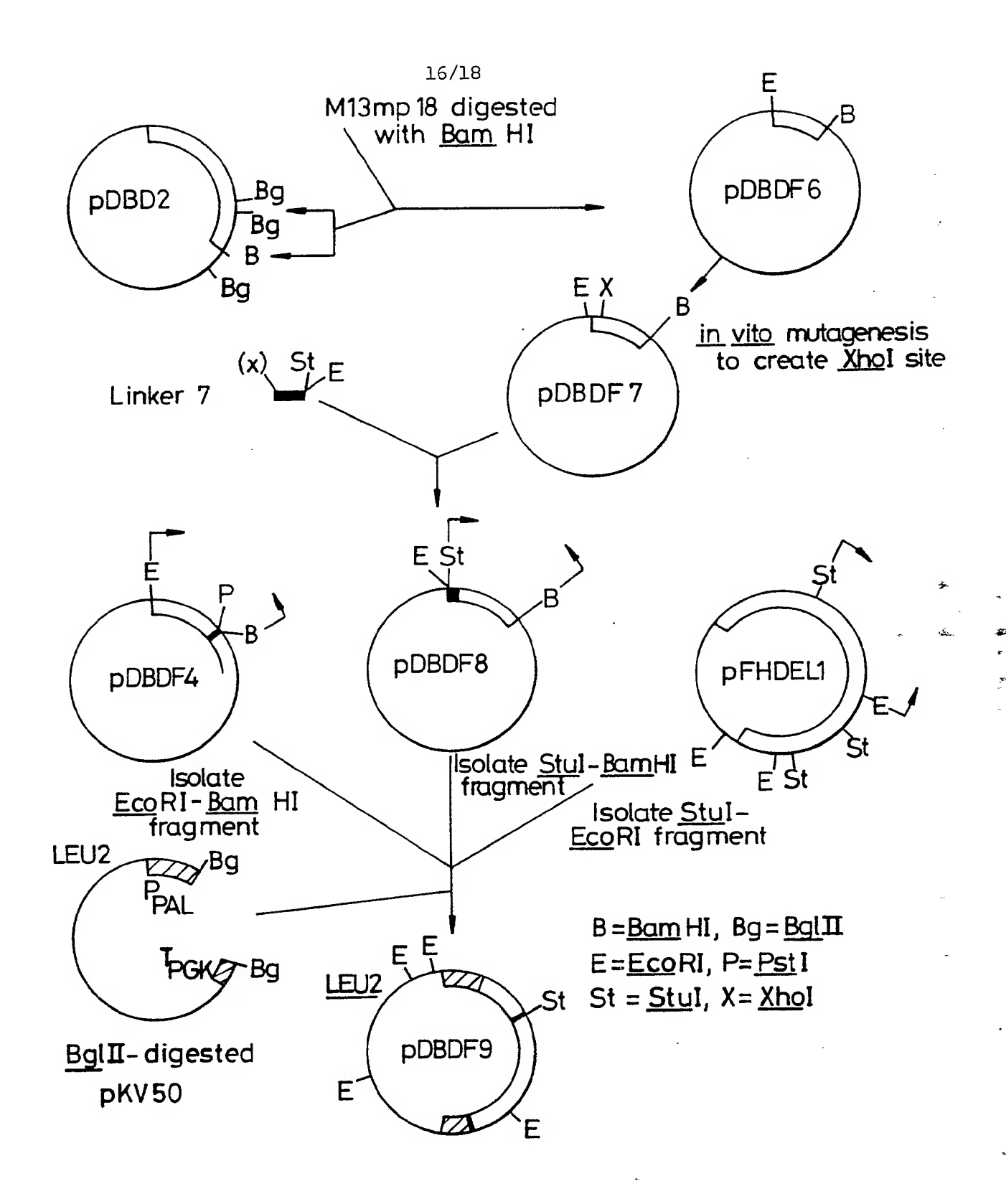
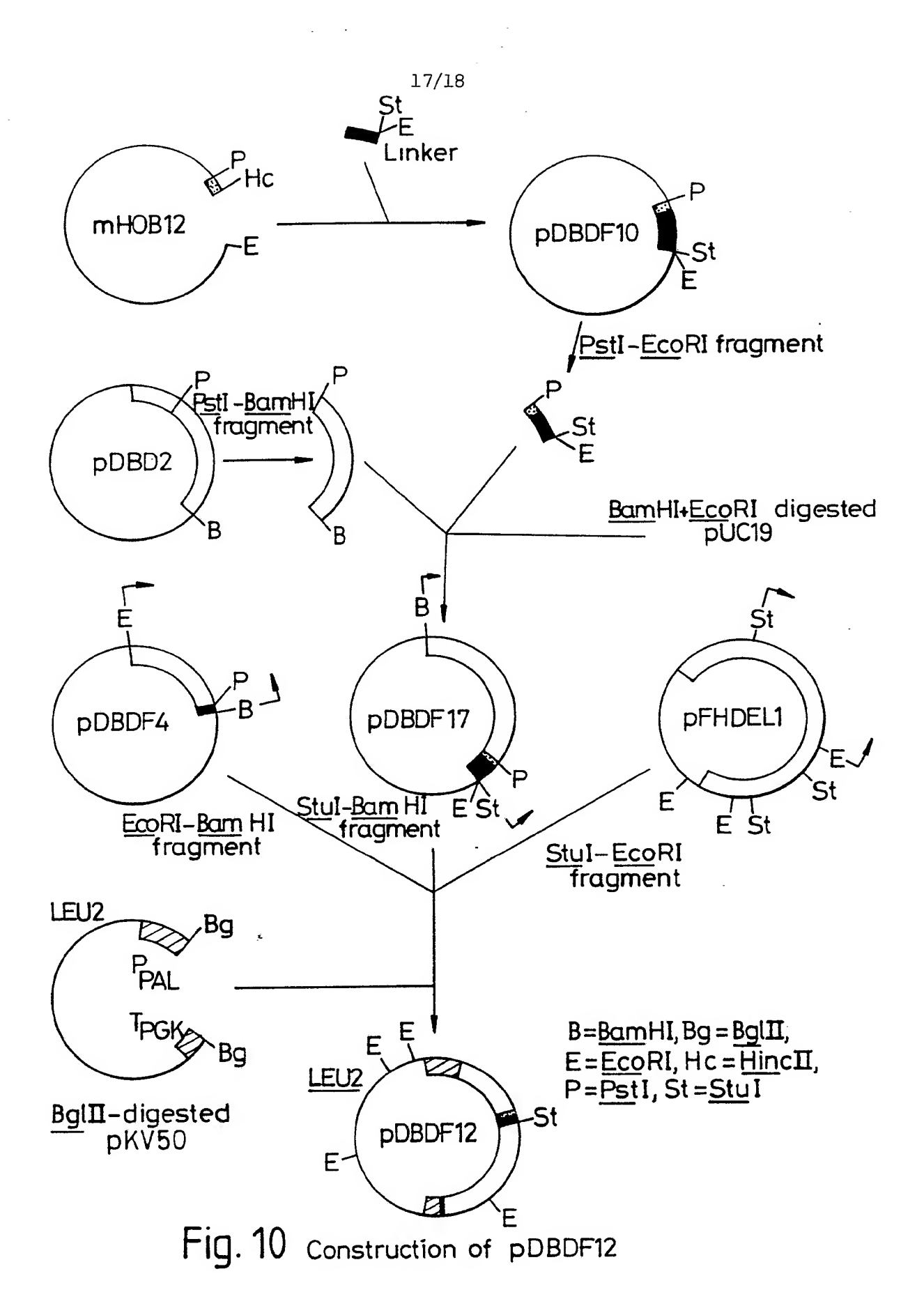


Fig. 9 Construction of pDBDF9

PCT/GB90/00650



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Figure 11

Name:

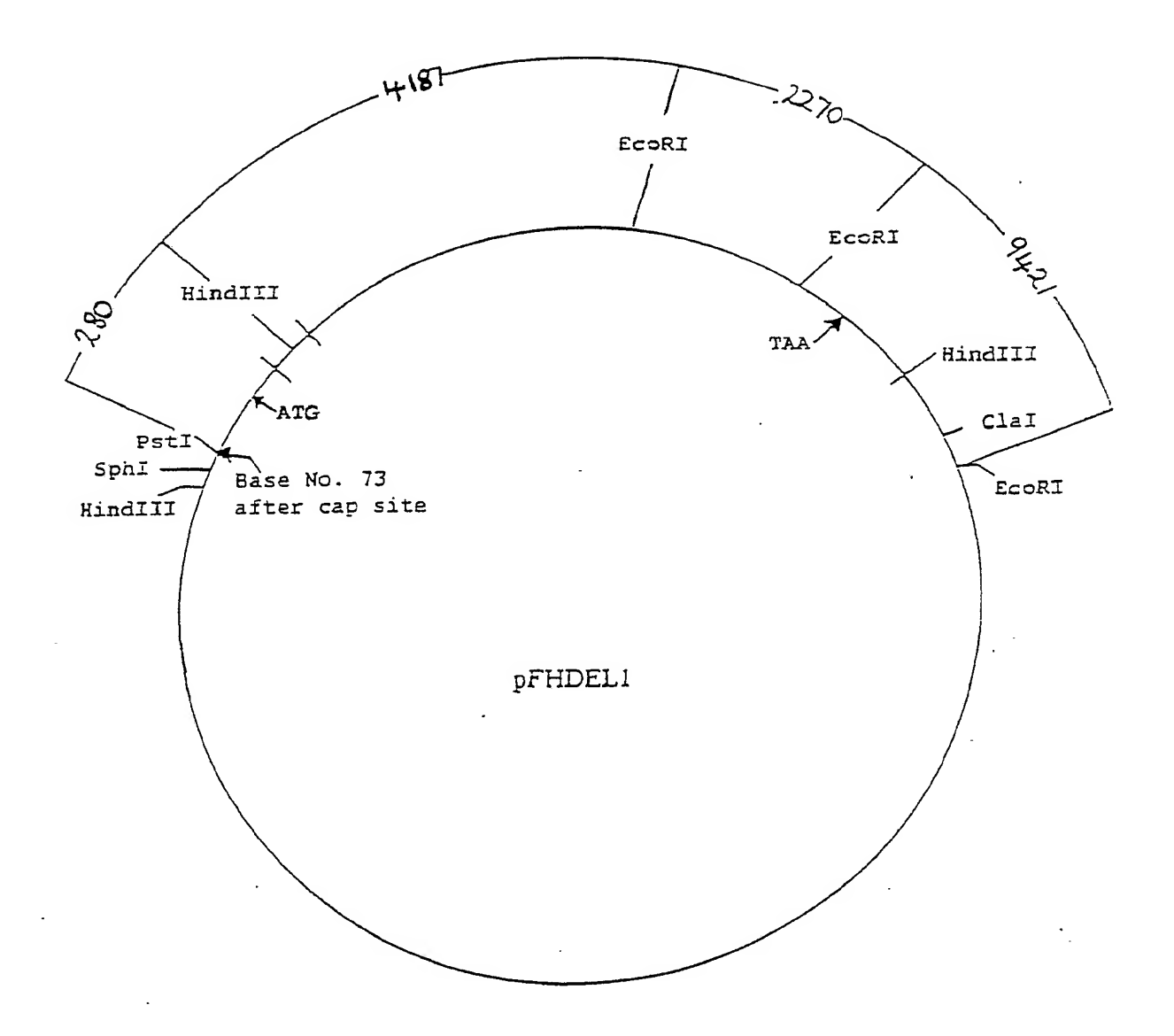
pFHDEL1

Yector:

pUC18 Ampfy 2860bp

Insert:

hFNcDNA - 7630bp



INTERNATIONAL SEARCH REPORT

International Application No PCT/GB 90/00650

I. CLAS	SIFICATION OF SUBJECT MATTER (if several classification	On symbols apply indicate all) 6					
Accordin	g to international Patent Classification (IPC) or to both National	Classification and IPC					
IPC ⁵	C 12 N 15/62, C 07 K 13/00						
II FIFLD	S SEARCHED						
Minimum Documentation Searched 7							
Classification System Classification Symbols							
E		incation Symbols					
IPC ⁵	C 12 N, C 12 P, C 07 K						
	Documentation Searched other than Note to the Extent that such Documents are In	finimum Documentation ncluded in the Fields Searched					
III DOCI	INTERCONSTRUCTOR OF STRUCTURE						
Category *	Citation of Document, 11 with Indication, where appropria						
		te, or the relevant passages 14	Relevant to Claim No. 13				
A	EP, A, 0308381 (SKANDIGEN e 22 March 1989	et al.)	-				
T	EP, A, 0322094 (DELTA BIOTE 28 June 1989 (cited in the application)	CHNOLOGY LTD)					
"A" documents of the Anti-	ment defining the general state of the art which is not idered to be of particular relevance or document but published on or after the international date of another date. The ment which may throw doubts on priority claim(s) or is cited to establish the publication date of another on or other special reason (as specified) The ment referring to an oral disclosure, use, exhibition or means Then the priority date claimed THE CATION	later document published after the or priority date and not in conflict cited to understand the principle invention document of particular relevance cannot be considered novel or clinvolve an inventive step document of particular relevance cannot be considered to involve an document is combined with one of ments, such combination being obtin the art. document member of the same particular relevance of Mailing of this International Search CO. 18, 90	t with the application but or theory underlying the claimed invention annot be considered to the claimed invention inventive step when the remove other such docuvious to a person skilled tent family				
nternational	Searching Authority Signa	iture of Authorized Officer M.	SOTELO				
	EUROPEAN PATENT OFFICE	IM					

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

GB 9000650

SA 36670

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 31/07/90

Patent document cited in search report	Publication date 22-03-89	Patent family member(s)		Publication date 17-07-89 17-04-89 15-03-89 23-03-89
EP-A- 0308381		SE-B- 459586 AU-A- 2420488 SE-A- 8703539 WO-A- 8902467		
EP-A- 0322094	28-06-89	AU-A-	2404688	18-05-89
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